WHAT IS CLAIMED IS

1. A phenoxypropylamine compound of the formula (I)

wherein each symbol in the formula means as follows:

- 5 a bond represented by a solid line and a dotted line shows a double bond or a single bond;
 - X is a hydrogen atom, a hydroxy group, a C_1-C_8 alkoxy group, an acyloxy group or an oxo group;

provided that when R^1 is a group of the following formula (2),

- 10 X should not be a hydrogen atom;
 - R^1 is a group of the following formula

$$-N \xrightarrow{Ar} -N \xrightarrow{N-z-R^2},$$

$$(1) \qquad (2) \qquad (2)$$

$$-N \xrightarrow{Z-R^5} \qquad -N \xrightarrow{Z-R^5}$$

$$(3) \qquad or \qquad (4)$$

wherein

y is 0 or S,

15 Ar is optionally substituted aromatic hydrocarbon,

 ${\ensuremath{\mathsf{R}}}^2$ is optionally substituted aryl group or optionally substituted aromatic heterocyclic group,

 R^5 is optionally substituted aryl group or optionally substituted aromatic heterocyclic group,

 20 Z is void or $-CH_2-$, and

 R^6 is hydrogen atom, hydroxy group, acetamido group,

 R^3 is a hydrogen atom, a $C_1 - C_{18}$ alkyl group or a halogen atom;

- - E ******

V is $-CH_2-$, -O-, -S- or the formula $-N(R^4)-$ wherein R^4 is hydrogen atom, C_1-C_{18} alkyl group or optionally substituted aralkyl group;

W is void or $-CH_2-$ or -C(=0)-;

is a C_1-C_4 hydroxyalkyl group, an acyl group, an optionally substituted saturated or unsaturated heterocyclic group, an optionally substituted fused heterocyclic group, a C_1-C_4 alkylsulfonyl group or the formula $-Q-R^9$

10 wherein

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Q is -C(=0)-, -C(=S)-, $-CH_2$ - or $-S(=0)_2$ -, and R⁹ is a group of the following formula

or -NH-NH-R¹⁵

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wherein R^{10} and R^{11} are each independently hydrogen atom, C_1-C_{18} alkyl group, optionally substituted aryl group, optionally substituted aralkyl group or alkoxy group, R^{12} is hydrogen atom, optionally

atom, phenyl group, C_1-C_4 alkyl group, C_1-C_2 halogenated alkyl group, halogen atom, C_2-C_4

alkenyl group, C_1 - C_4 hydroxyalkyl group, alkoxyalkyl group, alkyloxycarbonyl group, optionally substituted amino group, acetamido group, carboxyl group, acyl group, optionally substituted alkyloxy group, alkylthio group or cyano group;

provided that when R^1 is a group of the above formula (2), R^7 should not be C_1-C_4 hydroxyalkyl group or acyl group, and R^{10} and R^{11} are not each hydrogen atom at the same time; or

 $\ensuremath{\mbox{R}^{7}}$ and $\ensuremath{\mbox{W}}$ in combination may form a ring of the following formula

wherein

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E is oxygen atom or sulfur atom, and

Q' is an optionally substituted 4 to 7-membered heterocycle having 1 or 2 hetero atom(s) selected from the group consisting of nitrogen atom and oxygen atom in the ring, in which case V is hydrogen atom; and

Ra, Rb and Rc are each independently a hydrogen atom, a C_1-C_{18} alkyl group, a hydroxy group, a C_1-C_8 alkoxy group, a halogen atom, an acyl group, a nitro group or an amino group;

provided that when R^7 and W are bonded to form a ring of the above formula (14), Ra, Rb and Rc are not each hydroxy group or C_1-C_8 alkoxy group;

an optically active compound thereof, a pharmaceutically

2. The compound of claim 1, which is represented by the formula (I)

wherein each symbol in the formula means as follows:

- is a hydrogen atom, a hydroxy group, a C_1-C_8 alkoxy group, an acyloxy group or an oxo group;
 - R¹ is a group of the following formula

$$-N \longrightarrow_{Y} -N \longrightarrow_{N-Z-R^2}$$

$$-N \longrightarrow_{Z-R^5} -N \longrightarrow_{Z-R^5}$$

$$(3) \qquad \text{or} \qquad (4)$$

wherein

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10 Y is 0 or S,

Ar is optionally substituted benzene or naphthalene,

R² is optionally substituted aryl group or optionally substituted aromatic heterocyclic group,

R⁵ is optionally substituted aryl group or optionally substituted aromatic heterocyclic group,

Z is void or $-CH_2-$, and

 R^6 is hydrogen atom, hydroxy group, acetamido group, carboxyl group, alkoxycarbonyl group, cyano group or C_1-C_8 alkoxy group;

20 \mbox{R}^3 is a hydrogen atom, a $\mbox{C}_1\mbox{-}\mbox{C}_{18}$ alkyl group or a halogen atom;

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optionarry substituted alarkyr group;

25 W is void or $-CH_2-$ or -C(=0)-;

 R^7 is a C_1-C_4 hydroxyalkyl group, an acyl group, an optionally substituted saturated or unsaturated heterocyclic group, an optionally substituted fused heterocyclic group, a C_1-C_4 alkylsulfonyl group or the formula $-O-R^9$

wherein

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Q is -C(=0), -C(=S), $-CH_2$ - or $-S(=0)_2$ -, and R⁹ is a group of the following formula

or -NH-NH-R¹⁵

wherein R^{10} and R^{11} are each independently hydrogen atom, C_1 - C_{18} alkyl group, optionally substituted aryl group, optionally substituted aralkyl group or alkoxy group, R^{12} is hydrogen atom, optionally substituted aryl group, C_1 - C_{18} alkyl group, C_1 - C_8 alkoxy group or acyl group, and R^{15} is hydrogen atom, phenyl group, C_1 - C_4 alkyl group, C_1 - C_2

arkoxyalkyr group, alkyloxycarbonyr group, optionally substituted amino group, acetamido

group, carboxyl group, acyl group, optionally substituted alkyloxy group, alkylthio group or cyano group; and

Ra, Rb and Rc are each independently a hydrogen atom, a C_1-C_{18} alkyl group, a hydroxy group, a C_1-C_8 alkoxy group, a halogen atom, an acyl group, a nitro group or an amino group;

provided that when R^1 is a group of the above formula (2), R^7 should not be C_1 - C_4 hydroxyalkyl group or acyl group, and R^{10} and R^{11} are not each hydrogen atom at the same time; an optically active compound thereof, a pharmaceutically acceptable salt thereof or a hydrate thereof.

- 3. The compound of claim 2, which is represented by the formula (I) wherein each symbol in the formula means as follows:

 - X is a hydroxy group;
- 20 R¹ is a group of the following formula

$$-N \longrightarrow_{\mathbb{R}^6}^{\mathbb{Z}-\mathbb{R}^5} \qquad -N \longrightarrow_{\mathbb{Z}-\mathbb{R}^5} \mathbb{Z}-\mathbb{R}^5$$

wherein

- is optionally substituted phenyl group or naphthyl group,
- Z is void, and R^6 is hydrogen atom;
 - R^3 is a hydrogen atom or a C_1-C_4 alkyl group;
 - V is $-CH_2-$, -O-, -S- or $-N(R^4)-$
 - w is void;
 - R^7 is a group of the following formula

or the formula $-CO-R^9$ wherein

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 R^8 is hydrogen atom, phenyl group, C_1 - C_4 alkyl group, C_1 - C_2 halogenated alkyl group, halogen atom, C_2 - C_4 alkenyl group, C_1 - C_4 hydroxyalkyl group, alkoxyalkyl group, alkyloxycarbonyl group, optionally substituted amino group, acetamido group, carboxyl group, acyl group, optionally substituted alkyloxy group, alkylthio group or cyano group, and is a group of the following formula

$$R^{10}$$
 R^{11}
 R^{11}
 R^{12}
 R^{12}
 R^{12}
 R^{12}
 R^{12}
 R^{13}
 R^{14}
 R^{15}
 R^{15}

(9)

wherein R^{10} and R^{11} are each independently hydrogen atom, C_1-C_{18} alkyl group, optionally substituted aryl group, optionally substituted aralkyl group or alkoxy group, and R^{12} is hydrogen atom, optionally substituted aryl group, C_1-C_{18} alkyl group, C_1-C_8 alkoxy group or acyl group; and

or

(10)

Ra, Rb and Rc are each a hydrogen atom;

(8)

4. The compound of claim 2 or claim 3, which is represented by

the formula (I')

wherein each symbol is as in claim 2, an optically active compound thereof, a pharmaceutically 5 acceptable salt thereof or a hydrate thereof.

- 5. The compound of claim 2, which is selected from the group consisting of
- (1) 1-(4-(2-hydroxy-3-(4-(naphthalen-2-y1)piperidino) -
- 10 propyloxy) benzo (b) furan-2-ylcarbonyl) pyrrolidine,
 - (2) 4-(4-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)-propyloxy)benzo(b)furan-2-ylcarbonyl)morpholine,
 - (4) 4-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)propyloxy)-N,N-dimethylbenzo(b)furan-2-carboxamide,
- 15 (12) 1-(4-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)-propyloxy)benzo(b)thiophen-2-ylcarbonyl)pyrrolidine,
 - (13) 4-(4-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)-propyloxy)benzo(b)thiophen-2-ylcarbonyl)morpholine,
 - (15) 4-(2-hydroxy-3-(4-(naphthalen-1-yl)piperidino)propyloxy)-
- 20 N, N-dimethylbenzo (b) thiophene-2-carboxamide,
 - (17) 4-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)propyloxy)-N,N-dimethylbenzo(b)thiophene-2-carboxamide,
 - (20) 4-(7-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)-propyloxy)benzo(b)furan-2-ylcarbonyl)morpholine,
- 25 (21) 7-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)propyloxy)-N,N-dimethylbenzo(b)furan-2-carboxamide,

(30) = (2 hydroxy 3 (4 (maphinalen 2 yl)piperidino)propyloxy)

30 N,N-dimethyl-1-methylindole-2-carboxamide,

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- (35) 1-(2-(5-methyl-1,2,4-oxadiazol-3-yl)benzo(b)furan-4-
- yloxy)-3-(4-(naphthalen-2-yl)piperidino)-2-propanol,
- (37) 1-(2-(5-methyl-1,3,4-oxadiazol-2-yl)benzo(b)furan-4-yloxy)-3-(4-(naphthalen-2-yl)piperidino)-2-propanol,
- $_{5}$ (38) 1-(2-(5-trifluoromethyl-1,3,4-oxadiazol-2
 - yl) benzo(b) furan-4-yloxy)-3-(4-(naphthalen-2-yl)piperidino)-2-propanol,
 - (39) 1-(2-(5-methyl-1,3,4-oxadiazol-2-yl)benzo(b)furan-7-yloxy)-3-(4-(naphthalen-2-yl)piperidino)-2-propanol,
- 10 (42) 1-(2-(5-methyl-1,3,4-oxadiazol-2-yl)-1H-indole-4-yloxy)-3-(4-(naphthalen-2-yl)piperidino)-2-propanol,
 - (44) 1-(2-(3-methyl-1,2,4-oxadiazol-5-yl)benzo(b)furan-4-yloxy) 3-(4-(naphthālēn-2-yl)piperidino)-2-propanol,
 - (48) 1-(2-(5-methyloxazol-2-yl)benzo(b)furan-7-yloxy)-3-(4-
- 15 (naphthalen-2-yl)piperidino)-2-propanol,
 - (81) 3-(4-(3,4-dichlorophenyl)piperidino)-1-(2-(5-methyloxazol-2-yl)benzo(b) furan-4-yloxy)-2-propanol,
 - (88) 1-(4-(3,4-dichlorophenyl)piperidino)-3-(2-(5-methyl-1,3,4-oxadiazol-2-yl)benzo(b) furan-4-yloxy)-2-propanol, and
- 20 (93) 3-(4-(3,4-dimethylphenyl)piperidino)-1-(2-(5-ethyl-1,3,4-oxadiazol-2-yl)benzo(b)furan-4-yloxy)-2-propanol, an optically active compound thereof, a pharmaceutically acceptable salt thereof or a hydrate thereof.
- 25 6. The compound of claim 1, which is represented by the formula (I)

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double bond of a single bond,

X is a hydrogen atom, a hydroxy group, a C_1-C_8 alkoxy

group or an acyloxy group;

R¹ is a group of the following formula

$$-N$$
 $N-Z-R^2$ $-N$ $Z-R^5$ $-N$ $Z-R^5$ (4)

wherein

- is optionally substituted aryl group or optionally substituted aromatic heterocyclic group,
 - R⁵ is optionally substituted aryl group or optionally substituted aromatic heterocyclic group,
 - Z is void or $-CH_2-$, and
- 10 R^6 is hydrogen atom, hydroxy group or C_1-C_8 alkoxy group;
 - \mathbb{R}^3 is a hydrogen atom, a C_1-C_{18} alkyl group or a halogen atom;
 - ${\ensuremath{\mathsf{R}}^7}$ and ${\ensuremath{\mathsf{W}}}$ are bonded to form a ring of the following formula

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wherein

- E is an oxygen atom or a sulfur atom, and
- Q' is an optionally substituted 4 to 7-membered heterocycle having 1 or 2 hetero atom(s) selected from the group consisting of nitrogen atom and oxygen atom in the ring,

and V is hydrogen atom; and

Ra, Rb and Rc are each independently a hydrogen atom, a C_1-C_{18} alkyl group, a halogen atom, an acyl group, a nitro group or an amino group;

an optically active compound thereof, a pharmaceutically

30 formula (I) wherein each symbol in the formula means as

follows:

a group of the following formula

is a group of the following formula

wherein

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E is an oxygen atom or a sulfur atom,

q is 0, 1, 2 or 3,

 $R^{4'}$, $R^{7'}$ and $R^{8'}$ are each independently a hydrogen atom, a C_1-C_{18} alkyl group, an optionally substituted aryl group or an optionally substituted aralkyl group, and other symbols are as defined in claim 6, an optically active compound thereof, a pharmaceutically acceptable salt thereof or a hydrate thereof.

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8. The compound of claim 6, which is represented by the formula (I) wherein each symbol in the formula means as follows:

a bond represented by a solid line and a dotted line shows a double bond;

X is a hydroxy group;

R¹ is a group of the following formula

$$-N$$
 $Z-R^5$
 $-N$
 $Z-R^5$

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 R^5 is optionally substituted phenyl group or naphthyl

group,

Z is void, and

R⁶ is hydrogen atom;

 R^3 is a hydrogen atom or a C_1-C_4 alkyl group;

5 a group of the following formula

is a group of the following formula

$$(CH2) q R4' (19')$$

wherein q is 1 and $R^{4'}$ is hydrogen atom or C_1-C_4 alkyl group; and

Ra, Rb and Rc are each a hydrogen atom; an optically active compound thereof, a pharmaceutically acceptable salt thereof or a hydrate thereof.

9. The compound of claim 6, which is represented by the formula (I")

wherein each symbol is as as defined in claim 6, an optically active compound thereof, a pharmaceutically acceptable salt thereof or a hydrate thereof.

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(306) 5-(2'-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)-

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propyloxy) benzylidene) -1,3-dimethylimidazolidine-2,4-dione,
         (307) \alpha-(2'-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)-
        propyloxy) benzylidene) -γ-butyrolactone,
         (308) \alpha-(2'-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)-
  5 propyloxy) benzylidene) -γ-butyrolactone,
         (309) \alpha-(2'-(2-hydroxy-3-(4-(naphthalen-2-yl))piperidino)-
        propyloxy) benzylidene) -γ-butyrolactone,
         (310) \alpha-(2'-(3-(4-(3-fluoro-4-methylphenyl)piperidino)-2-
        hydroxypropyloxy) benzylidene) -y-butyrolactone,
10 (311) \alpha - (2' - (3 - (4 - (3, 4 - dimethylphenyl))))) = 2 - (311) \alpha - (2' - (3 - (4 - (3, 4 - dimethylphenyl))))
        hydroxypropyloxy) benzylidene) -y-butyrolactone,
         (312) \alpha-(2'-(3-(4-chloro-3-fluorophenyl)piperidino)-2-
        hydroxypropyloxy) benzylidene) -y-butyrolactone,
        (313) \alpha-(2'-(3-(4-(4-chloro-3-trifluoromethylphenyl)-
15 piperidino) -2-hydroxypropyloxy) benzylidene) -γ-butyrolactone,
        (314) \alpha-(2'-(2-hydroxy-3-(4-(naphthalen-1-yl)piperidino)-
       propyloxy) benzylidene) -γ-butyrolactone,
        (315) \alpha-(2'-(2-hydroxy-3-(4-(naphthalen-2-yl))piperidino)-
       propyloxy) benzylidene) -\delta-valerolactone,
20 (316) \alpha-(2'-(2-hydroxy-3-(4-(naphthalen-2-yl)piperidino)-
       propyloxy) benzylidene) -γ-valerolactone,
        (319) 3-(2'-(2-hydroxy-3-(4-(naphthalen-2-yl))piperidino)-
       propyloxy) benzylidene) -2-pyrrolidone,
        (322) 3-(2'-(2-hydroxy-3-(4-(naphthalen-2-y1)piperidino)-
25 propyloxy)benzylidene)-1-methyl-2-pyrrolidone, and
        (325) \alpha - (2' - (2-hydroxy-3 - (4 - (6-methoxynaphthalen-2 - (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) + (325) +
       yl)piperidino)propyloxy)benzylidene)-γ-butyrolactone,
       an optically active compound thereof, a pharmaceutically
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11 A pharmagnitical agent depositions on the same

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acceptable salt thereof or a hydrate thereof.

- 12. The pharmaceutical agent of claim 11, which is an agent for the treatment of depression.
- 13. A pharmaceutical composition comprising at least one member selected from the group consisting of a compound of claim 1, an optically active compound thereof, a pharmaceutically acceptable salt thereof and a hydrate thereof, and a pharmaceutically acceptable carrier.
- 10 14. The pharmaceutical composition of claim 13, which is an agent for the treatment of depression.
- 15. A 5HT_{IA} antagonist having a selective serotonin reuptake inhibitory action, which comprises a compound of claim 1, an optically active compound thereof, a pharmaceutically acceptable salt thereof or a hydrate thereof.
- 16. A selective serotonin reuptake inhibitor having a $5\mathrm{HT}_{1A}$ antagonistic action, which comprises a compound of claim 1, an optically active compound thereof, a pharmaceutically acceptable salt thereof or a hydrate thereof.
 - 17. A compound of the formula (II)

25 wherein each symbol in the formula means as follows:

y is a hydrogen atom a hydroxy grown of a live.

... a group or six rosrowing folimera

$$-N \xrightarrow{Ar} -N \xrightarrow{Z-R^5} -N \xrightarrow{Z-R^5} Z^{-R^5}$$
(1) (3) (4)

wherein

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Y is O or S,

5 Ar is optionally substituted benzene or naphthalene,

R² is optionally substituted aryl group or optionally substituted aromatic heterocyclic group,

R⁵ is optionally substituted aryl group or optionally substituted aromatic heterocyclic group,

10 \underline{Z} is void or $-CH_2$, and

 R^6 is hydrogen atom, hydroxy group, acetamido group, carboxyl group, alkoxycarbonyl group, cyano group or $C_1\text{-}C_8$ alkoxy group,

provided that when V is $-N(R^4)-$, R^6 should not be hydroxy group;

 R^3 is a hydrogen atom, a C_1-C_{18} alkyl group or a halogen atom;

V is $-CH_2-$, -O-, -S- or the formula $-N(R^4)-$ wherein

20 R^4 is hydrogen atom, C_1-C_{18} alkyl group or optionally substituted aralkyl group;

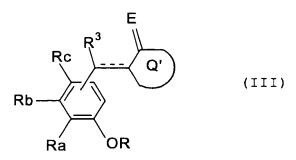
W is void, $-CH_2-$ or -C(=0)-;

 R^{14} is a hydrogen atom or a C_1-C_4 alkyl; and

Ra, Rb and Rc are each independently a hydrogen atom, a C_1-C_{18} alkyl group, a hydroxy group, a C_1-C_8 alkoxy group, a halogen atom, an acyl group, a nitro group or an amino group;

an optically active compound thereof a pharmacautically

18. A compound of the formula (III)



wherein each symbol is as follows:

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R is an allyl group or a 2,3-epoxypropan-1-yl group; a bond represented by a solid line and a dotted line shows a double bond or a single bond;

E is an oxygen atom or a sulfur atom;

 R^3 is a hydrogen atom, a C_1-C_{18} alkyl group or a halogen atom;

Q' is an optionally substituted 4 to 7-membered

heterocycle having 1 or 2 hetero atom(s) selected from
the group consisting of nitrogen atom and oxygen atom
in the ring; and

Ra, Rb and Rc are each independently a hydrogen atom, a C_1-C_{18} alkyl group, a hydroxy group, a C_1-C_8 alkoxy group, a halogen atom, an acyl group, a nitro group or an amino group;

an optically active compound thereof, a pharmaceutically acceptable salt thereof or a hydrate thereof.

20 19. The compound of claim 18, wherein, in the formula (III), each symbol is as follows:
the group of the following formula

is a group of the following formula

wherein

E is oxygen atom or sulfur atom,

q is 0, 1, 2 or 3,

 $R^{4'}$, $R^{7'}$ and $R^{8'}$ are each independently hydrogen atom, C_1 — C_{18} alkyl group, optionally substituted aryl group or optionally substituted aralkyl group, and other symbols are as defined in claim 18, an optically active compound thereof, a pharmaceutically acceptable salt thereof or a hydrate thereof.

- 20. A compound selected from the group consisting of 2-(4-methoxybenzo(b)furan-2-yl)-5-methyl-1,3,4-oxadiazole, 2-(4-hydroxybenzo(b)furan-2-yl)-5-methyl-1,3,4-oxadiazole,
- (S) -2-(4-glycidyloxybenzo(b) furan-2-yl) -5-methyl-1,3,4oxadiazole,
 - 2-(7-methoxybenzo(b)furan-2-yl)-5-methyl-1,3,4-oxadiazole,
 - 2-(4-(methoxymethyloxy)benzo(b)thiophen-2-yl)-5-methyl-1,3,4-oxadiazole,
- 20 2-(4-hydroxybenzo(b) thiophen-2-yl)-5-methyl-1,3,4-oxadiazole,
 4-benzyloxy-2-(5-methyl-1,3,4-oxadiazol-2-yl)-1H-indole,
 2-(7-methoxybenzo(b) furan-2-yl)-5-phenyl-1,3,4-oxadiazole,
 2-(4-methoxybenzo(b) furan-2-yl)-5-trifluoromethyl-1,3,4-oxadiazole,
 oxadiazole,
- 25 2-(4-hydroxybenzo(b) furan-2-yl)-5-trifluoromethyl-1,3,4-

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2-(7-methoxybenzo(b)furan-2-yl)-5-trifluoromethyl-1,3,4-

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oxadiazole,
   2-(7-hydroxybenzo(b)furan-2-yl)-5-trifluoromethyl-1,3,4-
   oxadiazole,
    (S) -2-(7-glycidyloxybenzo(b) furan-2-yl)-5-trifluoromethyl-
 5 1,3,4-oxadiazole,
   N'-(4-methoxybenzo(b) furan-2-ylcarbonyl) propionohydrazide,
   2-(4-methoxybenzo(b)furan-2-yl)-5-ethyl-1,3,4-oxadiazole,
   2-(4-hydroxybenzo(b) furan-2-yl)-5-ethyl-1,3,4-oxadiazole,
   (S) -2-(4-glycidyloxybenzo(b) furan-2-yl)-5-ethyl-1,3,4-
10 oxadiazole,
   2-(4-methoxybenzo(b) furan-2-yl)-5-methyl-1,3,4-thiadiazole,
   2-(4-hydroxybenzo(b) furan-2-yl)-5-methyl-1,3,4-thiadiazole,
   (S) -2-(4-glycidyloxybenzo(b) furan-2-yl)-5-methyl-1,3,4-
   thiadiazole.
15 5-ethoxycarbonyl-2-(4-methoxybenzo(b)furan-2-yl)-1,3,4-
   oxadiazole,
   5-ethoxycarbonyl-2-(4-hydroxybenzo(b)furan-2-yl)-1,3,4-
   oxadiazole,
   5-(4-(methoxymethyloxy)benzo(b) furan-2-y1)-2,3-dihydro-1,3,4-
20 oxadiazole-2-thione,
   5-(4-(methoxymethyloxy) benzo(b) furan-2-yl)-2-methylthio-1,3,4-
   oxadiazole,
   5-(4-hydroxybenzo(b) furan-2-yl)-2-methylthio-1,3,4-oxadiazole,
   5-(4-(methoxymethyloxy)benzo(b)furan-2-y1)-2,3-dihydro-1,3,4-
25 oxadiazol-2-one,
   5-(4-(methoxymethyloxy)benzo(b)furan-2-yl)-2-methoxy-1,3,4-
   oxadiazole,
   (S) - 5 - (4 - glycidyloxybenzo(b) furan - 2 - yl) - 2 - methoxy - 1, 3, 4 - yl)
   oxadiazole,
2-\text{ethoxy}-5-(4-(\text{methoxymethyloxy})\text{benzo(b)}\text{furan}-2-\text{yl})-1,3,4-
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JKdUldIOle,

Aradiamola

2-(1-methylethyloxy)-5-(4-(methoxymethyloxy)benzo(b)furan-2-

yl)-1,3,4-oxadiazole and

5

(S) -2-(1-methylethyloxy)-5-(4-glycidyloxybenzo(b)furan-2-yl)-1,3,4-oxadiazole.